# , UK Patent Application

(19) **GB** 

(11) 2 251 186<sub>(13)</sub>A

(43) Date of A publication 01.07.1992

(21) Application No 9026278.3

(22) Date of filing 04.12.1990

(71) Applicants
Randall Neal Gatz
Chemin de la Montau, CH-1251 Genolier, Vaud,
Switzerland

Peter Alfred Bromley 37 Chemin Cret-de-la-Neige, CH-1234 Vessy, Geneva, Switzerland

(72) Inventors
Randall Neal Gatz
Peter Alfred Bromley

(74) Agent and/or Address for Service Murgitroyd & Company Mitchell House, 333 Bath Street, Glasgow, G2 4ER, United Kingdom (51) INT CL\* A61K 37/02 39/00

(52) UK CL (Edition K)

A5B BHA B170 B180 B31X B31Y B317 B823 B826
B829 B835 B838 B842
U1S S1332 S2416 S2419

(56) Documents cited
GB 2221157 A EP 0322990 A1 WO 88/10120 A1
WO 85/05034 A1
Chin.exp.limmunol. 1990. 81, 189-194
Autoimmunity 1990, 7, 237-244
Immunology 1969, 16(2), 157-165

(58) Field of search
UK CL (Edition K) A5B BHA
INT CL<sup>5</sup> A61K 37/02 39/00
Online databases: WPI, DIALOG/PHARM

## (54) Polypeptide for use in treatment of autoimmune disease

(57) The use of a polypeptide comprising an amino acid sequence not homologous to a sequence synthesised by the cells of the patient, for the manufacture of a medicament for the treatment of an autoimmune disease is described.



#### Autoimmune Disease Treatment 1 2 This invention relates to the treatment of autoimmune 3 diseases, and especially the prophylactic treatment of 4. such diseases. 5 6 Stress of a varied nature, induced as a result of heat shock, nutrient deprivation, oxygen radicals and other g. forms of metabolic disruption, including infection by 9 certain viruses, bacteria and protozoans, as well as 10 certain cases of cellular transformation, all lead to 11 the increased synthesis of a family of proteins 12 collectively known as stress proteins or heat shock 13 14 proteins. 15 stress proteins are among the most highly 16 conserved and abundant proteins found in nature. 17 Further these proteins have been shown to be among the 18 dominant antigens recognised in immune responses to a 19 broad spectrum of pathogens. A review of the 20 interrelationships between stress proteins, infection 21 and immune surveillance has recently appeared, in 22

which a clear analysis of these relationships is

24 25

23

provided (13).

It has become apparent in recent years that a 1 relationship exists between so-called stress or 2 shock proteins and certain immune responses to 3 infection and to the development of autoimmunity. 4 an example, the analysis of cell-mediated and humoral 、 5 responses to a variety of bacterial and parasitic 6 7 pathogens has shown that heat shock proteins are often. strongly immunogenic during infection (1-8). 8 9 10 Proteins involved in immune responses to certain parasitic diseases such as malaria, shistosomiasis, 11 leishmanisis, trypanosomiasis and filariasis, have 12 been identified as members of the hsp 70 and 90 gene 13 families. Further antigens related to hsp 70 and GroEL 14 families have been shown to play a role in T cell and 15 B cell recognition during bacterial infections 16 including leprosy, tuberculosis and Q. fever. 17 18 mycobacterial GroEL stress protein has been identified as the target of a T cell clone capable of causing 19 autoimmune disease in a rat model of adjuvant-induced 20 arthritis (9). Similar results have been obtained as 21 concerns the small heat shock proteins, since an 22 immunologically important 19 Kd protein antigen of 23 24 Mycobacterium leprae has been sequenced, and shown to 25 have considerable amino acid sequence homology to the 26 soybean 19Kd heat shock protein. 27 Elevated responses to the GroEL stress protein have 28 been found by testing T cells from synovial infiltrates 29 of rheumatoid arthritis patients (10). Autoantibodies 30 31 to hsp 90 have been reported in systemic lupus erythrematosus (SLE) (11). In addition, elevated 32 antibody responses to hsp70 and GroEL stress proteins 33 have been found in SLE and in rheumatoid arthritis 34

35

(12).

1 The stress proteins are remarkable in their 2 evolutionary conservation: hsp90, hsp70, and hsp60 3 proteins are found in all prokaryotes and eukaryotes. In fact comparison of almost any two hsp70 proteins from two different organisms indicates an amino acid 6 homology of around 50%. The major stress proteins 7 occur at low levels in normal, unstressed cells, but 8 accumulate to very high levels in cells undergoing 9 stress. A striking example is the case of E. coli 10 hsp60, which accounts for 1.6% of total cell protein 11 under normal growth conditions, and can accumulate to 12 15% of total cell protein after heat shock (14). 13 Stress proteins appear to fulfil vital roles in cells, 14 both in the absence and in the presence of stress. 15 They appear to be involved in the assembly and 16 disassembly of protein complexes, and hsp70 proteins 17 are important for the translocation of certain 18 proteins through cellular membranes (15). Stress 19 proteins appear to interact with many different 20 proteins, for example, hsp90 has been found to interact 21 with steroid hormone receptors and with viral and 22 cellular kinases. Hsp70 proteins bind to DNA 23 replication complexes, clathrin baskets, the cellular 24 tumour antigen p53, and immunoglobulin heavy chains. 25 Plant hsp60 interacts with Rubisco, which fixes CO2 in 26 chloroplasts, and may be the most abundant protein in 27 the biosphere (16). The interaction of stress 28 proteins with multiple proteins may provide an 29 explication for the evolutionary constraints imposed 30 31 on their amino acid sequences. 32 33 Stress proteins have an almost certain role in

protecting cells and organisms from the deleterious

effects of heat and other stresses.

1 It seems clear that the tight sequence regulation 2 imposed on many heat shock protein sequences throughout 3 evolution has led to such retained sequences between those of the host and those of the infectious agent 5 having a significant degree of identity. Clearly the 6 reaction of the host immune system against antigens of 7 the infecting organism could lead to the raising of antibodies against heat shock proteins. The sequence 9 homology within the heat shock protein family thus 10 points to conserved sub-sequences of heat shock 11 proteins as being serious candidates for inducing an 12 immune response that can have specificity against self 13 sequences, with the consequence of inducing an 14 autoimmune reaction and the associated disease states. 15 16 The reports referenced above indicate that stress 17 proteins, such as the heat shock proteins, provide 18 particularly attractive targets for immune recognition. 19 An analysis the cross reactivity of T cell responses to 2.0 stress proteins has been published recently (17), 21 wherein the presence of human T cells was demonstrated 22 that were capable of immune recognition of conserved 23 sequence determinants. These authors have proposed a 24 model in which immune responses to stress proteins 25 provide a link between infectious and autoimmune 26 diseases. 27 28 Although models of the role of stress proteins in 29 autoimmune diseases have been proposed, no-one has yet 30 suggested possible treatment for autoimmune diseases. 31 32 In accordance with a first aspect of the present 33 invention a method of treating an autoimmune disease in - 34 a patient comprises introducing a compound, comprising 35

an amino acid sequence of a protein which is not 1 homologous with amino acid sequences synthesised by 2 cells of the patient, into the patient. 3 4 In accordance with another aspect of the present 5 invention there is provided use of a compound 6 comprising an amino acid sequence of a protein for the 7 treatment of an autoimmune disease in a patient, 8 wherein the amino acid sequence is not homologous with 9 amino acid sequences synthesised by cells of the 10 11 patient. 12 Further, the invention provides a composition for 13 treatment of an autoimmune disease in a patient, 14 comprising a compound which comprises an amino acid 15 sequence of a protein which is not homologous with 16 amino acid sequences synthesised by cells of the 17 patient, in combination with a pharmaceutical carrier. 18 19 Still further, the invention provides the use of a 20 compound comprising an amino acid sequence of a protein 21 which is not homologous with amino acid sequences 22 synthesised by the cells of a patient for the 23 manufacture of a medicament for the treatment of an 24 autoimmune disease in the patient. 25 26 Preferably, the compound comprises a peptide which 27 comprises the amino acid sequence and typically, the 28 protein is a stress or heat shock protein. 29 Preferably, the treatment is prophylactic. 321 Typically, the compound could be introduced into a 33° patient by incorporation in a cream or ointment, in a

soluble glass, in slow release capsules, transdermal

30

31

patches, injected, or even administered orally or in 2 suppository form. 3 Preferably, the amino acid sequence has antigenic 4 5 properties. 6 7 The amino acid sequence could be naturally occurring or be synthesised. If the amino acid sequence is synthesised then the peptide could comprise a number of 9 different amino acid sequences and/or multiples of the 10 11 same amino acid sequence. 12 The invention described here is based on the 13 above-detailed conservation of heat shock sequences and 14 their implication in autoimmune diseases. Contrary to 15 the identity of certain conserved sequences, this 16 invention, is based on the hypervariable sequences of 17 18 proteins. Prior immunisation with natural or synthetic peptides representing such non-conserved, 19 variable or hypervariable stress protein sequences of 20 origin from infectious agents of bacterial and other 21 parasitic pathogens, induces antibody responses 22 against the stress proteins of the infecting organism, 23 24 and these specifically induced antibodies are incapable of recognising self stress protein sequences. 25 rapid recognition of infectious agent - specific stress 26 27 proteins by specific pre-existing antibodies raised against non-homologous peptides from invading stress 28 29 proteins should allow the elimination of these stress 30 proteins before they are able to elicit potentially 31 autoimmune responses. 32 This invention concerns the immune recognition of 33 peptide epitopes of specific heat shock or stress 34 proteins, and the development of peptide-based therapy 35

or prevention based on such epitopes. I 2 Examples of the invention will now be described. 3 4 1. Analysis of stress protein peptide sequences 5 6 In order to practice the preventive/therapeutic 7 approach described in this invention, it is necessary to examine in detail the amino acid sequences of human 9 heat shock proteins, and of those of organisms 10 infecting human beings with whom correlations of immune 11 12 diseases exist. 13 Our initial approach was to assemble a table of certain 14 of the known sequences of stress proteins from human 15 and infectious agent sources. A selection of these 16 sequences are presented in Appendix 1. A thorough 17 analysis of sequence homology between members of each 18 of the stress protein families indicates that for 19 each of the principle stress protein families, hsp70, 20 hsp90 and hsp27, certain sequences have been highly 21 conserved throughout evolution, whereas parts of the 22 stress proteins contain amino acid sequences that are 23 highly differentiated. One assumes that the 24 conservational pressures concerning the retained 25 sequences are associated with critical structural or 26 functional aspects of these important proteins. 27 variable regions are presumably of less critical 28 structural or functional importance, thus escaping 29 from the conservative pressure/selection activities 30 prevailing in evolving organisms. 31 32 33 2. Selection of candidate peptide vaccines 34

3.5

The selection of useful candidate peptides capable of 1 eliciting an immune response specifically against the stress proteins of the infectious agent is based on 3 two major criteria: 5 6 The non-identity of selected peptide sequences, and their lack of resemblance to highly, or partially 7 conserved stress protein sequences, common to human 8 and infectious agent proteins. 9 The selection of such non-conserved sequences is derived from a reverse 10 11 analysis of amino acid sequence homologies, in other 12 words, concentrating on the non-homologous sequences 13 evident from homology analyses such as those shown in 14 (1) and in appendix 2. 15 16 For a thorough selection of sequence differences 17 versus sequence homology, it is instructive to, in addition to amino acid identity, to look at 18 19 replacements by highly conserved amino acids. Examples of such substitutions are the following groups: 20 21 (aspartic acid and glutamic acid), (lysine and arginine), (serine and threonine), (phenylalanine and 22 tyrosine), and (isoleucine, leucine, valine and 23 24 methionine). 25 26 An analysis of the antigenic potential of selected 27 peptide sequences. Where information is available, 28 peptide epitopes that conform to the criteria of both points i) and ii), and which can be demonstrated to be 29 immunodominant, are preferred examples of the 30 preventive/therapeutic peptides described in this 31 32 invention. 33 34 Examples of the amino acid sequences of some selected 35 peptides that reply to the criteria of point i) are

presented in appendix 2. I Examples of group i) peptides that are expected to 3 have considerable immunogenic potential have been selected on the basis of presently accepted criteria 5 of immunological potential. Examples of certain 6 peptides with pronounced antigenicity are shown in 7 appendix 3. 8 9. Non-homologous sequence comparison of the known stress IO: protein and related antigen sequences from humans and 11 from infectious agents has been performed. In the case 12 of Plasmodium falciparum, in addition to regions of 13 extensive homology of amino acid sequence between the 14 two proteins, clear regions of extensive lack of 15. homology are also detectable, and the following 16 sequence fragments, depicted using the one and 17 three-letter amino acid abbreviations derived from the 18 IUPAC-IUB Commission on Biochemical Nomenclature (see 19 Table 1), illustrate this example:-20 21 ALIGNMENT OF RESIDUES 133 TO 254 OF 75KDa antigen of  $\underline{P}$ 22 Falciparum TO RESIDUES 357 TO 635 OF HSP70 HUMAN 23 24 ENYCYGVKSSLEDKIKEKLQPAEIETCMKTITTILEWLEKNQLAGKDEYE 25 ----- KNALES-Y-AFNMKSA- VEDEG LKGKIS-E 26 27 AKQKEAESVCAPIMSKIY-QDAA-GAAGGMPGGM-P-GGMPGGMP GGMNF 28 ADKKKVLDKCQEVIS- WLDANTLA EKDEFEHKRKELEQVCNPIISGL-Y 29 30 PG-GMPG-AGMPGNAP---AGSGPTVEEVV 31 QGAGGPGPGGFGAQGPKGGSGSGPT----32. 33 Examples of non-homologous peptides are shown in bold 34 letters. The second peptide of HSP70 human shown in

;	bold above, denoted "Peptide example 1", has been
2	- compared to the sequence of the corresponding and
:	of Mycobacterium tuberculosis and its highly unique
4	sequence has little or no counterpart in the sequence
5	of tubercular origin.
. 6	
7	TESTAMENT OF RESIDUES 8 TO 11 OF PERTIPE 1 TO DESTRUCE
8	1 TO 127 OF 71KDa antigen M.tuberculosis
9	
10	<u>KRK</u> <u>E</u>
11	KEDIDRMIKDAEAHAEEDRKRREEADVRNGAETLVYNTEKFVKEQREGG
12	
13	Clearly other peptide sequences unique to an infectious
14	agent antigen exist and will have value in the
15	applications described in this invention. In and and
16	identity such sequences, extensive cloping comments
17	and sequence analysis of infectious agent anti-
18	will be required. Such research, although tochnically
19	diddods, is quite within the realms of evicting
20	technology. Similarly, once new sequences are
21	established, the presence or absence of amino acid
22	sequence nomologies can be determined either
23	visually, or through the use of any number of amatous
24 25	or commercial sequence analysis software programs our
26	intention here is to demonstrate the general procedure
27	for identifying, and applying both specific
28	non-homologous and specific homologous stress and
29	infectious agent antigen peptide seguences to the
30	vaccination, therapeutic and cosmetic applications
31	described herein.
32	
33	
	7 The Detical
35	3 The Rational Design of Synthetic Peptides
J J	

This invention is not limited to naturally occurring variant sequences within stress proteins, nor is it limited to the selection and use of a single variant epitope. For example, synthetic peptides could be used. In addition, the peptide could be synthesised to 5 have combinations of different variant sequences or multiples of variant sequences. By synthesising peptides comprising different variant sequences and/or multiples of the same variant sequence it may be 9 possible to design peptides having a stronger immune 10 response against stress proteins of infectious 11 organisms but which do not recognise human stress 12 epitopes. 13 14 A recent analysis of variant peptide epitopes of myelin basis protein (MBP), and their influence on the

15 16 incidence of experimental autoimmune encephalomyelitis 17 (EAE) has indicated that synthetic variants of an 18 N-terminal MBP peptide can have greatly altered 19 properties of binding to cell surface glycoproteins 20 encoded by the major histocompatibility complex (MHC) 21 (18). In other words, the efficacy of the complex 22 interactions associated with the elicitation of an 23 effective immune response against peptide antigens, can 24 be altered and improved in some cases, by the use of 25 synthetic variants of natural antigens. The subject 26 of this invention comprises those variant peptide 27 sequence approaches that are taught by the authors 28

29

of reference 18, amongst others.

30

An efficient mapping procedure for identifying protein 31 antigenic determinants has been described that would 32 be of use in the selection of useful antigenic

33 determinants for the applications taught in this 34

invention (19). Clearly classical chemical, enzymatic 35

and combined synthetic procedures can be utilised to produce candidate peptides, once identified and 2 selected, for the vaccine applications described here. 3 A naturally expected limitation of the peptide vaccines 5 that can be produced using this described procedure derives from the fact that about one third of monoclonal and polyclonal antibodies made by immunising with native protein react with assembled topographic sites (20). These assembled determinants 9 10 may not form the appropriate structure outside of a proteins native environment. This limitation is not 11 expected to significantly limit the practical use of 12 this invention. 13 14 Studies concerning T Cell recognition and activation 15 have indicated that it may be possible to design 16 peptides with predictable and advantageous properties 17 These authors have described two approaches for 18 immunomodulation that could be useful for the design 19 of therapeutic strategies against autoimmune 20 encephalomyelitis. The first approach consists of a 21 thorough molecular characterisation of an 22 encephalitogenic epitope, and the subsequent design of 23 peptide analogs that retain normal or increased major 24 25 histocompatibility complex binding properties, and that fail to activate disease-inducing T cells. Secondly, 26 novel properties of a heterocyclic peptide have been 27 described, with the result that the peptide is highly 28 antigenic in vitro, while being non-immunogenic in 29 These authors have been able to demonstrate the 30 vivo. feasibility of immune intervention in an immune disease 31 through the use of a synthetic peptide. These results 32 are complementary to the procedure we describe here, 33 but are not identical, nor do they in any way predict 34

the approach that we describe.

1 4 Applications of the stress protein peptides described 2 3 herein The basic tenant that we have developed herein is based 5 on the multiple observations that certain infectious 6 agent antigens are closely related in amino acid 7 sequence to human stress proteins, and that immune 8 reactions against such antigens can cross react with 9 the human proteins, leading to the possibility of 10 developing autoimmune disease. Our invention describes 11 the selection of stress protein peptide sequences from 12 infectious agent antigens related to human stress 13 proteins, but which have little or no sequence homology 14 within such human stress proteins. The injection of 15 such non-homologous peptides into human beings, for 16 instance in an emulsification with Freunds complete 17 adjuvant, would provide a route of effective 18 vaccination against subsequent autoimmune disease 19 20 induced as mentioned above. The antibodies raised through such vaccination are specific to the selected 21 infectious agent antigen from which the vaccinating 22 peptide was derived. Such induced antibodies are 23 specific to infectious agent antigens, thus explaining 24 their efficacy in the application of this invention. 25 26 Further, since the vaccinating agent is a small 27 peptide, instead of a large, complex protein such as 28 human factor VIII, or factor IX, it is not compulsory 29 30 to use an injection as a means of delivering the peptide to a human subject. We thus reserve in our 31 application the administration of the kinds of peptides 32 described by transdermal applications, a number of 33 which are presently commercialised with considerable 34 35 success.

Further still, since certain major diseases that are thought to have their origin in autoimmune diseases, such as arthritis and rheumatism, the peptides of this invention can be applied externally, in both local and cosmetic application to painful joints and articulations resulting from these prevalent diseases. For example, the peptides could be administered to a patient by incorporation in a cream or ointment, in a soluable glass, in slow release capsules, transdermal patches, injected, or even administered orally or in suppository form. In addition, due to the nature of amino acid sequences it is unlikely that treatment using these substances will produce the unpleasant side effects which are normally associates with drugs. 

•)

I			APPENDIX 1		
2					
3	NO	N-HOMOLOGOUS	S SEQUENCES V	WHICH ARE ALS	SO .
4			GENICS ARE I		
5		UNDERLINING	AND NON-HOMO	LOGOUS ONLY	
6	SI			OLD LETTERIN	iG
7		•			
8					
9		SEQUENCE OF	HUMAN STRES	S PROTEINS	
10					
11					
12	A) Sequenc	e HSP90 Huma	n .		
13					
14	Rebbe N F,	Ware J, Ber	tina M, Modr	ich P, Staff	ord D 1
15		5-245(1987)			
16	EMBL; M166	60; HSHSP90			
17	KW Heat Sh	ock. Sequenc	e 724 AA; 83	293 MW	
18					
19					-
20					
21	MPEEVHHGEE	EVETFAFQAE	IAQLMSLIIN	TFYSNKEIFL	40
22	RELISNASDA	LDKIRYESLT			
23	ERTLTLVDTG	IGMTKADLIN	NLGTIAKSGT		
24	ADISMIGQFG	VGFYSAYLVA	EKVVVIRKHN	DDEQYAWESS	160
25	AGGSFTVRAD	HGEPIGMGTK	VILHLKEDQT	EYLEERRVKE	200
26	VVKKHSQFIG	YPITLYLEKE	REKEISDDEA	EEEKGEKEEE	240
27	DKDDEEKPKI	EDVGSDEEDD	SGKDKKKKTK	KIKEKYIDQE	280
28	ELNKTKPIWT	RNPDDITQEE	YGEFYKSLTN	DWEDHLAVKH	320
29	FSVEGQLEFR	ALLFIPRRAP	FDLFENKKKK	NNIKLYVRRV	360
30	FIMDSCDELI	PEYLNFIRGV	VDSEDLPLNI	SREMLQQSKI	400
31	LKVIRKNIVK	KCLELFSELA	EDKENYKKFY	EAFSKNLKLG	440
32	IHEDSTNRRR	LSELLRYHTS	QSGDEMTSLS	EYVSRMKETQ	480
33	KSIYYITGES	KEQVANSAFV	ERVRKRGFEV	VYMTEPIDEY	520
34	CVQQLKEFDG	KSLVSVTKEG	LELPEDEEEK	KKMEESKAKF	560
35	ENLCKLMKEI	LDKKVEKVTI	SNRLVSSPCC	IVTSTYGWTA	600

```
LRDNSTMGYM MAKKHLEINP
                                           DHPIVETLRQ
                                                       640
 1
     NMERIMKAQA
 2
     KAEADKNDKA
                  VKDLVVLLFE
                              TALLSSGFSL
                                           EDPQTHSNRI
                                                       680
     TYMIKLGLGI DEDEVAAEEP
                              NAAVPDEIPP
                                           LEGDEDASRM
                                                       720
                                                       724
     EEVD
 5
 6
 7
 8
     B) Sequence HSP70 Human
 9
     [1] Hunt C, Morimoto R I;
10
     Proc Natl Acad Sci USA 82:6455-6459(1985)
11
     EMBL; M11236; HSHSP701
12
13
     EMBL; MII717; HSHSP70D
     KW Heat Shock
14
     Sequence 640AA; 69867 MW
15
16
17
     MAKAAAVGID
                  LGTTYSCVGV
                              FQHGKVEIIA
                                          NDQGNRTTPS
                                                       40
18
     YVAFTDTERL
                  IGDAAKNQVA
                              LNPQNTVFDA KRLIGRKFGD
                                                       80
19
     PVVQSDMKHW
                  PFQVINDGDK
                              PKVQVSYKGE TKAFYPEEIS
                                                       120
                                                       160
20
     SMVLTKMKEI
                  AEAYLGYPVT
                              NAVITVPAYF
                                          NDSQRQATKD
                              IAYGLDRTGK GERNVLIFDL
                                                       200
21
     AGVIAGLNVL .
                 RIINEPTAAA
     GGGTFDVSIL
                  TIDDGIFEVK
                              ATAGDTHLGG
                                          <u>EDF</u>DNRLVNH
                                                       240 (3)
22
                                                       280
     FVEEFKRKHK
                  KDISQNKRAV
                              RRLRTACERF
                                          EGIDFYTSIT
23
     RARFEELAKR
                  TLSSSTOASL
                              EIDSLCSDLF
                                          RSTLEPVEKA
                                                       320(4)
24
                                                       360
                  IHDLVLVGGS
                              TRIPKVOKLL
                                          ODFFNGRDLN
25
     LRDAKLDKAQ
                                                       400
26
     KSINPDEAVG
                  YGAAVQAAIL
                              MGDKSENVQD
                                          LLLLDVAPLS
                                                       440
27
     LGLETAGGVM
                  TALIKRNSTI
                              PTKQTQIFTT
                                          YSDNQPGVLI
                              LSGIPPAPGV
                                          PQIEVTFDID
                                                       480 (1)
     QVYEGERAMT
                  KDNNLLGRFE
28
                              TITNDKGRLS
                                          KEEIERMVQE
                                                       520
29
     ANGILNVTAT
                 DKSTGKANKI
30
     AEKYKAEDEV
                 <u>ORERVSAKNĀ</u>
                              LESYAFNMKS
                                          AVEDEGLKGK
                                                       560 (<u>2</u>)
31
     ISEADKKKVL
                 DKCQEVISWL
                              DANTLAEKDE
                                          FEHKRKELEQ
                                                       600
                              FGAQGPKGGS
                                          GSGPTIEEVD
                                                       640
32
     VCNPIISGLY
                 QGAGGPGPGG
33
34
```

**.** ..

```
C) Sequence Human HSP27
   I
   Z
      Hickey E, Brandon S E, Potter R, Stein G, Stein J,
   3
      Weber L A;
   4
      Nucl. Acids Res 14:4127-4145(1986)
  5.
      EMBL;X03900; HSHSP27
  6
  7
      KW: HEAT SHOCK
      SEQUENCE 199 AA; 22327 MW;
  ä٠
  9
      MTERRVPFSL LRGPSWDPFR DWYPHSRLFD QAFGLPRLPE 40
 IO.
      EWSQWLGGSS WPGYVRPLPP AAIESPAVAA PAYSRALSRQ 80
 II
      LSSGVSEIRH TADRWRVSLD VNHFAPDELT VKTKDGVVEI
 12
                                                      120
      TGKHEERQDE HGYISRCFTR
 13
                             KYTLPPGVDP TQVSSSLSPE
                                                      160
 14:
      GTLTVEAPMP KLATQSNEIT IPVTFESRAQ LGGRSCKIR
                                                      200
 15.
 16
      D) Sequence Human HSP60
 17
 18
     Sequence not yet available, submitted for publication:
     Gupta R S, Jinal S, Harley C B and Dudani A K(1989)
 19
20
21
22.
                     SEQUENCE OF HSP60 YEAST
23.
24
25 -
     Reading D S, Hallberg R L and Myers A M (1989). Nature
26.
     <u>337</u> 655
27
     MLRSSVVRSR ATLRPLLRRA YSSHKILKFG VIGRASLLKG
28
                                                     40
29.
     VETLAIAVAA TLGPKGRNVL IEQPFGPPKI
                                         TKDGVTVAKS
                                                     80
30
     IVLKDKFINM GAKLLQIVAS KTNIAAGDGT
                                         TSATVLGRAI
                                                     120
     FTISVKNVAA GCNPMDLRRG SQVAVIKVIL
31
                                         FLSANKKEIT
                                                     160
     TSEEIAQVAT ISANGDSHVG KLLASAMEKV
32
                                         GKEGVITIRE
                                                     200
33
     GRITLEDELE VTEGMRFDRG FISPYFITDP
                                         KSSKVEFEKP
                                                     240
34
    LLLLSEKKIS SIQDILPALE ISNQSRRPLL IIAEDVDGEA
                                                     280
35
     LAACILNKLR GQVKVCAVKA PGFGDNRKNT
                                         IGDIAVLTGG
                                                     320
```

```
TVFTEELDLK PEQCTIENLG SCDSITVTKE DTVILNGSGP
  1
                                                     360
     KEAIQERIEQ IKGSIDITTT NSYEKEKLQE RLAKLSGGVA 400
  2
     VIRVGGASEV EVGEKKDRYD DALNATRAAV EEGILPGGGT 440
  3
     ALVKASRVLD EVVVDNFDQK LGVDIIRKAI TRPAKQIIEN 480
     AGEEGSVIIG KLIDEYGDDF AKGYDASKSE YTDMLATGII
                                                     520
 6
     DPFKVVRSGL VDASGVASLL ATTEVAIVDA PEPPAAAGAG 560
 7
     GMPGGMPG
                 MPGMM
                                                     600
 8
                 SEQUENCES OF BACTERIAL ANTIGENS
10
11
12
     A) Mycobacterium leprae
13
14
     18 KDa Antigen
15
16
     Nerland A H, Mustapha A S, Sweetser D, Godal T, Young R
17
     J Bacteriol 170 5919-5921 (1988)
18
     Sequence 148 AA; 16643MW;
19
20
     MLMRTDPFRE LDRFAEQVLG TSARPAVMPM
                                        DAWREGEEFV 40
21
     VGFDLPGKA
                 DSLDIDIERD VVTVRAERPG
                                        VDPDREMLAA 79
22
     ERPRGVFNRQ LVLGENLDTE RILASYQEGV
                                        LKLSIPVAER 119
23
     AKPRKISVDR GNNGHQTINK TPHEIIDA
24
25
26
     65 KDa Antigen
27
28
29
    Mehra V, Sweetser D and Young R A (1986) Proc Natl Acad
30
     Sci USA <u>83</u> 7013
31
32
    AA 589, MW 61,831
33
    The Underling Amino Acids Correspond To Antigenic
34
    Peptides.
```

```
40
     VPGRDGETQP ASCGRPSRAL HPASVSNGGC RSPVILASFL
 1
                                                    80
     IRRNHFAMAK TIAYDEEARR GLERGLNSLA DAVKVTLGPK
 2
                                                    120
     GRNVVLEKKW GAPTITNDGV SIAKEIELED PYEKIGAELV
 3
                                                    160
    KEVAKKTDDV AGDGTTTATV LAQALVKEGL RNVAAGANPL
 4
 5
     GLKRGIEKAV DKVTETLLKD AKEVETKEQI AATAAISAGD
                                                    200
 6
    QSIGDLIAEA MDKVGNEGVI TVEEESNTFG LQLELTEGMR
                                                    240
                                                    280
    FDKGYISGYF VIDAERQEAV LEEPYILLVS SKVSTVKDLL
 7
 8
    PLLEKVIQAG KSLLIIAEDV EGEALSTLVV NKIRGTFKSV
                                                    320
     AVKAPGFGDR RKAMLQDMAI LTGAQVISEE VGLTLENTDL
                                                    360
 9
     SLLGKARKVV MTKDETTIVE GAGDTDAIAG RVAQIRTEIE
                                                    400
10
    NSDSDYDREK LQERLAKLAG GVAVIKAGAA TEVELKERKH
11
                                                    440
                                        ALDKLKLTGD
                                                    480
12
    REIDAVRNAK AAVEEGIVAG GGVTLLQAAP
                                                    520
     EATGANIVKV ALEAPLKQIA FNSGMEPGVV AEKVRNLSVG
13
                                                    560
    HGLNAATGEY EDLLKAGVAD PVKVTRSALO
                                        NAASIAGLFL
14
                                                    600
15
     TTEAVVADKP EKTAAPASDP TGGMGGMDF
16
17
18
     70 KDa Antigen
19
20
    Not yet sequenced. Immunological cross-reactivity with
21
    the 71 KDa antigen of Mycobacterium tuberculosis (YOUNG
22
    ET AL Proc Natl Acad Sci USA 85, 4267-4270 (1988).
23
24
25
        Mycobacterium tuberculosis
26
    B)
27
28
    65 KDa Antigen
29
30
31
     Schinnick, T S (1987). Journal of Bacteriology 169
32
33
    1080
    AA 562, MW 59083
34
35
```

```
1
      RGCRHPVTPP
                  VSSPIRRNHF AMAKTIAYDE EARRGLERGL
                                                     40
  2
      NALADAVKVT
                  LGPKGRNVVL EKKWGAPTIT NDGVSIAKEI
                                                     80
  3
      ELETPYEKIG
                  AELVKEVAKK TDDVAGDGTT
                                         TATVLAQALV
                                                     120
      REGLRNVAAG ANPLGLKRGI EKAVEAKVTET LLKGAKEVET
  4
                                                     160
  5
      KEQIAATAAI
                  SAGDQSIGDL IAEAMDKVGN EGVITVEESN
                                                     200
      TFGLQLELTE GMRFDKGYIS GYFVTDPERQ EAVLEDPYIL
  6
                                                     240
  7
      LVSSKVSTVK
                  DLLPLLEKVI
                             GAGKPLLIIA EDVEGEALST
                                                     280
     LVVNKIRGTF
  8
                  KSVAVKAPGF
                             GDRRKAMLQD MAILTGGQVI
                                                     320
 9
      SEEVGLTLEN
                  ADLSLLGKAR
                             KVVVTKDETT IVEGAGDTDA
                                                     360
 10
     IAGRVAQIRQ
                  EIENSDSDYD REKLQERLAK LAGGVAVIKA
                                                     400
 11
     GAATEVELKE
                             NAKAAVEEGI VAGGGVTLLK
                  RKHRIEDAVR
                                                     440
12
     AAPTLDELKL
                 EGDEATGANI
                             VKVALEAPLK QIAFNSGLEP
                                                     480
13
     GVVAEKVRNL PAGHGLNAQT
                             GVYEDLLAAG VADPVKVTRS
                                                     520
14
     ALQNAASAIG LFLTTEAVVA
                             DKPEKEKASV
                                        PGGGDMGGMD
                                                     560
15
     F
                                                     600
16
17
18
     71 KDa Antigen
19
20
21
     Partial sequence, contains only the homolgy domain with
22
     HSP70
23
24
     Young D, Lathigra R, Hendrix R, Sweetser D, Young R,
25
     Proc Acad Sci
     USA 85, 4265-4270 (1988).
26
27
28
     EFQPSVQIQV YQGEREIAAH NKLLGSFELT
                                        GIPPAPRGIP
                                                    40
                                                        (1)
     QIEVTFDIDA NGIVHVTAKD KGTGKENTIR
29
                                        IQEGSGLSKE
                                                    80
30
     DIDRMIKDAE A<u>HAEEDRKRR EEADVRNGA</u>E
                                        TLVYNTEKFV
                                                    120 3,4
    KEQREGGSKV PEDTWRIGYF GHQVGDGEAG
31
                                        PGVAGSGASD
                                                    160 (2)
    LRSSSGCVTG HWRCPPRAAA GRCPPRLGM
32
                                                    200
33
34
```

```
C) Plasmodium falciparum (MALARIA)
  1
  2
  3
      90 KDa Antigen
  4
  5
  6
  7
     M Jendoubi, S Bonnefoy, Nucl Acids Res 16, 10928 (1988)
      Partial sequence, contains only the region of homology
     with HSP90
  9
 10
     KDFDGKKLKC CTKEGLDIHH SEEAKKDFET
11
                                         VIKDVLHKKV
                                                      40
     EKVVVCQRIT DSPCVLVTSE FGWSANMERI MKAQALRDNS 80
12
13
     MTSYMLSKKI
                 MEINARHPII
                             SALKQKADAD
                                                      120
                                         KSDKTVKYLI
14
     WLLFDTSLLT
                             TFSKRIHRMI
                 SGFFALEEPT
                                         KLGLSIDEEE
                                                     160
     NNDIDLPPLE
15
                 ETVDATDSKM
                             EEVD
                                                      200
16
17
18
     75 KDa Antigen
19
20
21
     Ardeshir F, Flint J E, Richman S and Reese R T, Embo J.
22
     6, 493-499
23
     (1987).
24
     Partial sequence from the first AA
25
26
     MLKLIERNTT IPAKKSQIFT TYADNQPGVL
                                         IQVYEGERAL
                                                     40
27
     TKDNNLLGKF HLDGIPFAPR KVPQIEVTFD
                                         IDANGILDVT
                                                     80
28
     AVEKSTGKQN HITITNDKGR LSQDEIDRMV
                                         NDAEKYLAED
                                                     120
     EENRKRIEAR NSLENYCYGV KSSLEDKIKE
29
                                         KLQPAEIETC
                                                     160
30
     MKTITTILEW LEKNQLAGKD EYEAKQKEAE
                                         SVCAPIMSKI
                                                     200
31
     YQDAAGAAGG MPGGMPGGMP
                             GGMPGGMNFP
                                         GGMPGAGMPG
                                                     240
32
     NAPAGSGPTV EEVVD
                                                     280
33
34
35
```

1	APPENDIX 2
2	
3	DIFFERENTIATION OF HOMOLOGOUS (UNDERLINE)
4	AND NON-HOMOLOUGOS SEQUENCES
5	
6	
7	A) Alignment of Residues 47 to 161 of partial sequence
8	F. Faiciparum 90KD to residues 581 to 699 of human
9	HSR90
10 11	
12	RI-DSPCVLVTSEFGWSANMERIMKAOALRDNSMTSYMLSKKIMEINAR
13	NRLVSSPCCIVTSTYGWTANMERIMKAQALRDNSTMGYMMAKKHLEINPD
14	UDITAL PROPERTY.
15	HPIISALK <u>OKADADKSDKTVKYL</u> IW <u>LLFDTSLLTSGFFALEEPTTFSKRI</u>
16	HPIVETLRQKAEADKNDKAVKDLVVLLFETALLSSG-FSLEDPQTHSNRI
17	UDVIVI or annual
18	HRMIKLGLSIDEEENN
19	YRMIKLGLGIDEDEVAAEE
20	
21	D. 33:
22	B) Alignment of residues 7 to 157 of partial sequence
23	of P. falciparum 70 KDa to residues 411 to 613 of human HSP70.
24	HSP/U.
25	· Main The same a second secon
26	NTTIPAKKSOIFTTYADNOPGVLIOVYEGERALTKDNNLLGKFHL
27	ALIKRNSTIPTKQTQIFTTYSDNQPGVLIQVYEGERAMTKDNNLLGRFEL
28	DCIDD) DDW DC
29	DGIPPAPRKVPOIEVTFDIDANGILDVTAVEKSTGKONHITITNDKGRLS
30	SGIPPAP-GVPQIEVTFDIDANGILNVTATDKSTGKANKITITNDKDRLS
31	QD <u>EIDRMV</u> ND <u>AEKYLAEDE</u> EN <u>RKRIEARNSLENY</u> CYGV <u>KS</u> SL <u>ED</u> K-I <u>K</u> E <u>K</u> LQ
32	KEEIERMVQEAEKYKAEDEVQRERVSAKNALESYAFNMKSAVEDEGLKGKIS
3 3	THE TALL THE
4	PAETCMKTITTILEWLEKNQLAGKDEYEAKQKEAESVCAPIMSKIYODA
5	EADKKKVLDKCQEVI-SWLDANTLAEKDEFEHKRKELEQVCNPIISGLYQGA

7	
I	
2	C) Alignment of residues 5 to 110 of M tuberculosis 71K to residues 430 to 548 of burns were
3	to residues 430 to 548 of human HSP70
4	
5	YSDNQPGVLIOVYEGERANTYRANDA
6	YSDNQPGVLIQVYEGERAMTKDNNLLGRFELSGIPPAP-GVPQIEVTFDI
7	·
8	DANGIVH <u>VTAKDKGTGKENTIRIQEGSG-LSKE</u> DID <u>RM</u> IKD <u>AE</u> AHAEEDR
9	DANGILNVTATDKSTGKANKITITNDKGRLSKEEIERMVQEAEKYKAEDE
10	THOMGRESKELLERMVQEAEKYKAEDE
11	KRREEADVRNGAE
12	VQRERVSAKNALESYAFNM
13	
14	·
15	
16	
17	$\cdot$
18	
19	
20	
21	
22	
23	
24	
25	
26	
27 -	
28	
29	
30	·
31	
32	

1	<u>APPENDIX 3</u>
2	
3	Antigenic Peptides of the 65 Kda Antigen of
4	Mycobacterium leprae
5	
6	MEHRA V, SWEETSER D and YOUNG R A (1986) Proc Natl Acad
7	Sci USA <u>83</u> 7013
8	
9	-NSLADAVKVTLGPKGRNVVLEKKWGAPTITNDGVS
10	-RNVAAGANPLGLKRGIEKAV
11	-ALDKLKLTGDEATGA
12	-GEYEDLLKAGVADP
13	-ASDPTGGMGGMDF
14	
15	<b>:</b>
16	
17	
18	
19	
20	·
21	
22	
23	
24	
25	
26	
27	
28	
29	
30	
31	
32	
33	
34	

1				
1				TABLE 1
2			•	
3		One and	Three Letter	Amino Acid Abbreviations
4				
5				•
6.		A	Ala	Alanine
7		C	Cys	Cysteine
8		D	Asp	Aspartic acid
9		· E	Glu	Glumatic acid
10		F	Phe	Phenylalanine
11		G	Gly	Glycine
12		H	His	Histidine
13		I	Ile	Isoleucine
14		K	Lys	Lysine
15		L	Leu	Leucine
16		М	Met	Methionine
17		N	Asn	Asparagine
18		P	Pro	Proline
19		Q	Gln	Glutamine
20		R	Arg	Arginine
21		S	Ser	Serine
22		T	The	Threonine
23 .		· <b>V</b>	Val	Valine
24		W	Trp	Tryptophane
25		Y	Tyr	Tyrosine
26		В	Asx	Asp or Asn (not
27				distinguished)
28		Z	Glx	Glu or Gln (not
29				distinguished)
30		X	x	Undetermined or atypical
31	-			amino acid
32		v		
33	From:	IUPAC-I	UB Commission	on Biochemical

33 From: IUPAC-IUB Commission on Biochemical

<sup>34</sup> Nomenclature, J Biol

<sup>35</sup> Chem 243, 3557-3559, 1968.

References

1

2 Young D.B, Lathigra R, Hendrix R, Sweetser D and 3 4 Young R A 1988. Stress proteins are immune targets in 5 leprosy and tuberculosis. Proc Natl Acad Sci USA <u>85</u> 4267. 6 7 Vodkin M H and Williams J C 1988. A heat shock 8 operon in Coxiella burnetti produces a major antigen 9 homologous to a protein in both mycobacteria and 10 Escherichia coli. J Bacteriol 170 1227. 11 12 Bianco A E, Favaloro J M, Burkot T R, Culvenor J 13 G, Crewther P E, Brown G V, Anders R F, Coppel R L and 14 Kemp D J 1986. A repetitive antigen of Plasmodium 15 falciparum that is homologous to heat shock protein 70 16 of Drosophila melanogaster. Proc Natl Acad Sci USA 83 17 18 8713. 19 Ardesshir F, Flint J E, Richman S J and Reese R T 20 1987. A 75Kd merozile surface protein of Plasmodium 21 falciparum which is related to the 70 Kd heat-shock 22 proteins. EMBO J 6 493. 23 24 Hedstrom R, Culpepper J, Harrison R A, Agabian N 25 and Newport G 1987. A major immunogen in Schistosoma 26 mansoni infections is homologous to the heat-shock 27 protein Hsp 70. J Exp Med 165 1430. 28 29 Selkirk M E, Rutherford P J, Denham D A, Partano F 30 and Maizels R M 1987. Cloned antigen genes of Brugia 31 filarial parasites. Biochem Soc Symp 53 91. 32 33 Dragon E A, Sias S R, Kato E A and Gabe J D 1987. 34 The genome of Trypanosoma cruzi contains a 35

\$1

```
constitutively expressed tandemly arranged multicopy
 1
     gene homologous to a major heat shock protein. Mol
 2
     Cell Biol 7 1271.
 3
 4
          Jendoub M and Bonneloy S 1988. Identification of
     a heat shock-like antigen in P. falciparum related to
     the heat shock protein 90 family. Nucleic Acids Res 16
     10928.
 9
          van Eden W, Thole J E R, van der Zee R, Noordzy A,
10
     van Embden J D A, Hensen E J and Cohen I R 1988.
11
     Coning of the mycobacterial epitope recognised by T
12
     lymphocytes in adjuvant arthritis. Nature 331 171.
13
14
          Res P C M, Schaar C G, Breedveld F C, van Eden W,
     10
15
     van Embden J D A, Cohen I R and de Vries R R P 1988.
16
     Synovial fluid T cell reactivity against 65 Kd heat
17
     shock protein of mycobacteria in early chronic
18
     arthritis. Lancet 478.
19
20
          Minota S, Koyasu S, Yahara and Winfield J 1988.
21
     11
     Autoantibodies to the heat shock protein hsp90 in
22
     systemic lupus erythrematosus. J Clin Invest 81 106.
23
24
          Tsoulfa G, Rook G A W, van Embden J D A, Young D
     12
25
     B, Mehlert A, Isenberg D A, Hay F C and Lydyard P M
26
            Raised serum IgG and IgA antibodies to
27
     mycobacterial antigens in rheumatoid arthritis. Annals
28
     of Rheumatic Diseases. 48 118.
29
30
          Young R A and Elliot T J 1989. Stress Proteins,
31
     Infection, and Immune Surveillance.
                                           Cell <u>59</u> 5
32
33
          Herendeen S L, van Bogelen R A and Neidhardt F C
34
     14
```

1979. J Bacteriol <u>139</u> 185.

```
1
  2
           Cheng M Y, Hartl F -U, Martin J, Pollock R A,
      15
      Kalousek F, Neupert W, Hallberg E M, Hallberg R L and
  3
      Horwich A L 1989. Nature 337 620.
  4
           Hemmingsen S M, Woolford C, van der Vies S M,
  6
      16
      Tilly K, Dennis D T, Georgopoulos C P, Hendrix R W and
  7
      Ellis R J 1988. Nature 333 330.
  8
  9
      17 Lamb J R, Bal V, Mendez-Samperio P, Mehlert A, So
 10
      A, Rothbard J, Jindal S, Young R A and Young D B 1989.
 11
      Stress proteins may provide a link between the immune
 12
      response to infection and autoimmunity. The Japanese
 13
      Society for Immunology 0953 8178/89, International
14
15
      Immunology Vol 1 No 2.
16
17
           Urban J L, Horvath S J and Hood L 1989.
     Autoimmune Recognition of Normal and Variant Peptide
18
     Epitopes and Peptide-Based Therapy. Cell 59 257.
19
20
          Mehra V, Sweetser D and Young R A 1986.
21
     mapping of protein antigenic determinants. Proc Natl
22
23
     Acad Sci USA 83 7013.
24
     20 Berzofsky J A 1985. Science <u>229</u> 932
25
26
          Wraith D C, Smilek D E, Mitchell D J, Steinman L
27
     21
     and McDevitt H O 1989. Cell <u>59</u> 247.
28
29
30
31
32
33
34
```

CLAIMS

2

4

1. A method of treating an autoimmune disease in a patient comprises introducing a compound, comprising an amino acid sequence of a protein which is not homologous with amino acid sequences synthesised by cells of the patient, into the patient.

11

12 2. Use of a compound comprising an amino acid
13 sequence of a protein for the treatment of an
14 autoimmune disease in a patient, wherein the amino
15 acid sequence is not homologous with amino acid
16 sequences synthesised by cells of the patient.

17

18 3. A composition for treatment of an autoimmune
19 disease in a patient, comprising a compound which
20 comprises an amino acid sequence of a protein
21 which is not homologous with amino acid sequences
22 synthesised by cells of the patient, in
23 combination with a pharmaceutical carrier.

24

25 4. The use of a compound comprising an amino acid 26 sequence of a protein which is not homologous with 27 amino acid sequences synthesised by the cells of a 28 patient for the manufacture of a medicament for 29 the treatment of an autoimmune disease in the 30 patient.

31

32

33 34

# Parents Act 1977 Examiner's report to the Comptroller under Section 17 (The Search Report)

# Application number

9026278.3

Section 1	Search Examiner
CI (Edition K ) A5B (BHA)	Search Examiner
at CI (Edition ) A61K 39/00, 37/02	C SHERRINGTON
Databases (see over)	Date of Search
ONLINE DATABASES: WPI, DIALOG/PHARM	3 FEBRUARY 1992

Documents considered relevant following a search in respect of claims

Category see over)	Identity of document and relevant passages	Relevant to claim(s)
X	GB 2221157 A (BIOGAL GYOGYSZERGYAR) especially page 1, line 12-14; Claim 19	4
х	EP 0322990 A1 (DE STAAT DER NEDERLANDEN) whole document	4
X	WO 88/10120 A1 (BRIGHAM AND WOMEN'S HOSPITAL) whole document especially page 6, line 19 - page 7, line 3; Example 6; Claims 1-10,13-19	4
A	WO 85/05034 A1 (UNIVERSITY OF LONDON ET AL) especially page 2, line 18 - page 3, line 4; Claims 3-5	4
x	Clin.exp.Immunal.1990,81,189-194 Prevention of adjuivant arthritis in rats by a nonapeptide from the 65-kd	<b>4</b>
x	Autoimmunity 1990,7,237-244  The immune response to Mycobacterial heat shock proteins	4
X	Immunology 1969,16(2),157-165 Inhibition of Adjuvant Arthritis by Protein Antigens	4

Category	Identity of document and relevant passages	Helevant to claim(s
Ì		•
	•	
	<b>'</b> .	

### Categories of documents

- X: Document indicating lack of novelty or of inventive step.
- Y: Document indicating lack of inventive step if combined with one or more other documents of the same category.
- A: Document indicating technological background and/or state of the art.
- P: Document published on or after the declared priority date but before the filing date of the present application.
- E: Patent document published on or after, but with priority date earlier than, the filing date of the present application.
- &: Member of the same patent family, corresponding document.

Databases: The UK Patent Office database comprises classified collections of GB, EP, WO and US patent specifications as outlined periodically in the Official Journal (Patents). The on-line databases considered for search are also listed periodically in the Official Journal (Patents).